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Barcoding bushmeat: molecular identification of Central African and South American harvested vertebrates

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Abstract The creation and use of a globally available database of DNA sequences from a standardized gene region has been proposed as a tool for species identification, assessing genetic diversity and monitoring the legal and illegal trade in wildlife species. Here, we contribute to the Barcode of Life Data System and test whether a short region of the mitochondrial cytochrome *c* oxidase subunit 1 (*COXI*) gene would reliably distinguish among a suite of commonly hunted African and South American mammal and reptile species. We used universal primers to generate reference barcode sequences of 645 bp for 23 species from five vertebrate families (*Crocodylidae*, *Alligatoridae*, *Bovidae*, *Suidae* and *Cercopithecidae*). Primer cocktails yielded high quality barcode sequences for 179 out of 204 samples (87.7%) from all species included in the study. For most taxa, we sequenced multiple individuals to estimate intraspecific

sequence variability and document fixed diagnostic characters for species identification. Polymorphism in the *COXI* fragment was generally low (mean = 0.24%), while differences between congeneric species averaged 9.77%. Both fixed character differences and tree-based maximum likelihood distance methods unambiguously identified unknown and misidentified samples with a high degree of certainty. Barcode sequences also differentiated among newly identified lineages of African crocodiles and identified unusually high levels of genetic diversity in one species of African duiker. DNA barcoding offers promise as an effective tool for monitoring poaching and commercial trade in endangered species, especially when investigating semi-processed or morphologically indistinguishable wildlife products. We discuss additional benefits of barcoding to ecology and conservation.

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Molecular forensics · Primates · Wildlife monitoring ·
Maximum likelihood phylogeny · Ungulates

Introduction

The hunting of tropical wildlife has historically been conducted for subsistence consumption and for local trade. But current trends in wildlife harvest from across the globe suggest that the volume of extraction of wild game, or “bushmeat”, has increased considerably, and many species are in sharp decline due to over exploitation (Albrechtsen et al. 2007; Bennett et al. 2007; Milner-Gulland et al. 2003; Redford 1992). Former locally-based subsistence economies have become global, and bushmeat is now a significant export product traded at regional and international

scales. Of equal importance to the deleterious impacts on wild populations and ecosystems promoted by unregulated harvest are the potential human and agricultural health impacts stemming from the increased potential for zoonotic disease transfer (Chomel et al. 2007; Milius 2005).

Considering the international trade in bushmeat and fisheries, estimates of its worth are in excess of US\$60 billion per year, with wildlife and wildlife products contributing US\$5–15 billion (Baker 2008). A significant portion of this trade is illegal (US\$5–8 billion, in Baker 2008), involving species that are protected by national laws and international conventions governing the use of wildlife and wildlife products. Given the illicit nature of the trade, it is difficult to accurately assess and monitor the volumes and species involved (Chomel et al. 2007; Milius 2005) and thus fully understand existing and potential impacts on economies, wildlife populations and health.

Existing legislation and treaties governing the trade in wildlife, such as the Convention on the International Trade of Endangered Species (CITES) and the United States Endangered Species Act (ESA), are based on the recognition of distinct population or taxonomic units. At a minimum, enforcement of regulations depends upon an ability to identify suspected illegal products at the species level. Accurate identification is often impaired due to the types of products involved, which are typically processed and difficult to identify using morphological techniques. To improve our ability to detect, monitor and control the trade in wildlife and wildlife products, more accurate and efficient methods of species identification are required. Identification of fish and wildlife species targeted for commercial trade is considered among the most useful applications of molecular ecology (Baker 2008).

The ability to identify wildlife products, whether as processed meat, skins or whole animals, is being formalized by the development of DNA sequence databases using a standardized gene fragment (Ratnasingham and Hebert 2007; Ross et al. 2003). A database of single-gene “barcodes” has been proposed to classify the complete diversity of life (Hebert et al. 2003a; Ratnasingham and Hebert 2007) and proponents argue that such a tool could variously be employed for defining taxonomic units for conservation (Neigel et al. 2007; but see Rubinoff 2006), biological inventory (Janzen et al. 2005) and species discovery (Bickford et al. 2007; Hebert et al. 2004; Kaila and Stahls 2006; Witt et al. 2006). The 5′ Folmer region of the mitochondrial (mtDNA) cytochrome *c* oxidase I (*COXI*) gene has been recommended as a standard for DNA barcoding (Folmer et al. 1994; Hebert et al. 2003a, b; Ivanova et al. 2007). Although there has been considerable criticism of the philosophical and practical underpinnings of DNA barcoding (DeSalle 2006; Fitzhugh 2006; Rubinoff 2006; Rubinoff et al. 2006;

Song et al. 2008), its application for species identification has largely been uncontested (e.g. Rubinoff et al. 2006).

The use of sequence data for investigations of endangered species collected in commercial markets is widely reported. While the *COXI* region has not been used as a standard metric, these studies have employed mitochondrial markers to identify samples to the species level. Yan et al. (2005) used the cytochrome *b* (*CYTB*) gene to identify Chinese alligators (*Alligator sinensis*) from fresh and partially cooked meat found in Chinese markets. Baker and colleagues used mtDNA sequences and microsatellites to identify endangered whale species sampled from markets in Korea and Japan (Baker et al. 1996, 2002). Using diagnostic characters in *CYTB*, Birstein et al. (1998) identified three species of caviar-producing sturgeon and discovered that nearly a quarter of commercially available caviar lots sold in New York City were mislabeled. Marko et al. (2004) also used *CYTB* sequences to determine that 77% of fish sold in the US labeled as red snapper were actually other species. Martin (1991) and Moura et al. (2008) used mtDNA sequences, including *COXI*, to identify commercially fished shark species when morphological characters (e.g. fins, heads) were equivocal in discerning among and within genera. The identification of endangered wildlife products in each of these cases would not have been possible without molecular methods because vendors had a vested interest in concealing the identity of the species being sold.

Ours is the first study to examine the utility of universal *COXI* primers as a standard metric to identify multiple species for monitoring the global trade in wildlife, with particular emphasis on species commonly traded in bushmeat markets. Our study includes species from five taxonomic families: bovids (duikers and spiral-horned antelope; genera: *Cephalophus*, *Tragelaphus*), suids (red river hog; genus: *Potamochoerus*), cercopithecoid primates (old world monkeys and mangabeys; genera: *Cercopithecus*, *Lophocebus*), alligators (genera: *Caiman*, *Melanosuchus*, *Paleosuchus*) and crocodiles (genera: *Crocodylus*, *Osteolaemus*, *Mecistops*) (Table 1). We obtained samples either from museum collections or from in situ captures. We use fixed diagnostic characters to compare interspecific and intergeneric levels of variation and bootstrap node support from a maximum likelihood (ML) tree-based approach to assess the monophyly of closely related species. Diagnostic characters and phylogenetic support were also used to assign a small number of unidentified samples to species and to examine the museum collection and field specimens for possible errors in species identification or labeling. The poor quality of some tissue used in this study reflects the suboptimal conditions encountered when working with material sampled from bushmeat markets, processed wildlife products, or reference museum samples

Table 1 Sample list, sample size, number of samples that failed to sequence, tissue type, sample origin, and conservation status for 11 genera included in the COX1 barcoding study. Unknown samples are listed with their respective known family classification

Order	Family	Species	Museum	Wild	Failed	Storage method	Origin	Status ^a
Crocodilia	Alligatoridae	<i>Paleosuchus palpebrosus</i>	4	–	2	Tissue (blood) in buffer	Brazil	Least concern
		<i>Paleosuchus trigonatus</i>	3	–	–	Tissue (blood) in buffer	Brazil	Least concern
		<i>Caiman crocodilus chiapasius</i>	4	–	–	Tissue (blood) in buffer, dried blood on gauze	Costa Rica	Least concern
		<i>Caiman yacare</i> ^b	8	–	3	Dried blood on gauze	Bolivia; Brazil; Peru	Least concern
		<i>Caiman c. crocodilus</i> × <i>yacare</i>	1	–	–	Tissue (blood) in buffer	Brazil	Least concern
		<i>Melanosuchus niger</i>	4	–	–	Dried tissue (non-blood)	Peru	Least concern
		<i>Caiman latirostris</i>	8	–	2	Tissue (blood) in buffer, dried blood on gauze	Bolivia; Brazil; Paraguay	Least concern
		Unknown caiman	1	–	–	Tissue (blood) in buffer, dried blood on gauze	Bolivia	Least concern
		<i>Crocodylus acutus</i>	1	–	–	Dried tissue (non-blood)	Oaxaca, Mexico	Vulnerable
		<i>Mecistops cataphractus</i>	–	10	–	Dried tissue (non-blood)	Gabon; Congo	Data deficient
Artiodactyl	Bovidae	<i>Crocodylus niloticus</i>	–	8	–	Dried tissue (non-blood)	Gabon	Least concern
		<i>Crocodylus niloticus</i>	–	1	–	Dried tissue (non-blood)	Congo	Least concern ^c
		<i>Osteolaemus tetraspis</i>	1	16	3	Dried tissue (non-blood)	Gabon; Cameroon	Vulnerable
		<i>Osteolaemus osborni</i>	–	11	1	Dried tissue (non-blood), tissue in buffer	Congo	Vulnerable
		Unknown African crocodile ^d	5	–	4	Dried tissue (processed leather)	Unknown	Vulnerable
		<i>Tragelaphus eurycerus eurycerus</i>	–	1	–	Dried blood on filter paper	Ndoki, Congo	Near threatened
		<i>Cephalophus nigrifrons</i>	–	5	–	Tissue in buffer	Ndoki, Congo	Least concern
		<i>Cephalophus leucogaster</i>	–	8	1	Tissue in buffer	Ndoki, Congo	Least concern
		<i>Cephalophus dorsalis</i> ^e	–	10	1	Tissue in buffer	Kabo, Congo	Least concern
		<i>Cephalophus callipygus</i>	–	26	3	Dried blood on filter paper	Ndoki, Congo	Least concern
Primates	Cercopithecidae	<i>Cephalophus monticola</i>	–	26	2	Dried blood on filter paper, tissue in buffer	Ndoki, Congo	Least concern
		Unknown ungulate	–	5	1	Dried blood on filter paper	Congo	Least concern
		<i>Potamochoerus porcus</i>	–	8	1	Tissue in buffer	Ndoki, Congo	Least concern
		<i>Cercopithecus cephus</i>	–	7	–	Dried blood on filter paper, tissue in buffer	Ndoki, Congo	Least concern
		<i>Cercopithecus nictitans</i>	–	8	1	Dried blood on filter paper	Ndoki, Congo	Least concern
		<i>Cercopithecus pogonias</i>	–	6	–	Dried blood on filter paper, tissue in buffer	Ndoki, Congo	Least concern
		<i>Lophocebus albigena</i>	–	8	–	Dried blood on filter paper	Ndoki, Congo	Least concern
		Total	40	164	25			

^a As established by the International Union for the Conservation of Nature's (IUCN) Red List of Threatened Species, www.iucnredlist.org

^b Includes one sample originally identified as *M. niger*

^c The IUCN has yet to recognize the new crocodile species and their status remains the same as the original taxa

^d From US Fish & Wildlife Service, National Wildlife Property Repository (http://www.fws.gov/mountain-prairie/law/property_repository.html)

^e Includes one sample originally identified as *C. callipygus*

and, therefore, provides a robust evaluation of the utility of DNA barcoding for wildlife monitoring and investigations.

Materials and methods

Species identification and sample collection

Specimens used in this study were identified and collected under four different scenarios. The first involved harvested mammals surveyed during a prior study of bushmeat hunting in the Republic of Congo (Table 1; Eaton 2002). Species identifications were made by MJE and trained field assistants using African mammal guides (Estes 1991; Kingdon 1997) in conjunction with the knowledge of local hunters. Voucher photographs were taken of all species and later confirmed using Nowak (1999). All samples, with the exception of five unidentified ungulates, were collected from freshly killed, whole animals in which species identification was unambiguous. The five unidentified samples were collected from partially butchered animals that could only be confirmed as belonging to ungulates based on hair pattern and coloration. Samples of Osborn's dwarf crocodiles (*Osteolaemus osborni*) and a single Nile crocodile (*Crocodylus niloticus*) were collected from intact, hunted animals during a subsequent study in the Republic of Congo (Eaton et al. 2009; Thorbjarnarson and Eaton 2004). The second sample collection scenario consisted of capture and identification of live African crocodiles for systematics and ecological research in the Republics of Congo and Gabon (Eaton 2006; Eaton and Barr 2005; Eaton et al. 2009; Thorbjarnarson and Eaton 2004). Although there have been recent revisions to the taxonomy of all African crocodiles (Eaton et al. 2009; Hekkala 2004; McAliley et al. 2006; Schmitz et al. 2003), identification to genus (and in most cases to species) was unambiguous. The third source of samples was voucher crocodilian specimens obtained from museum collections. These included dried blood and tissue from six species of South American caiman, the American crocodile (*Crocodylus acutus*) and the Central African dwarf crocodile (*Osteolaemus tetraspis*) provided by the Yale Peabody Museum (YPM) and the American Museum of Natural History (AMNH; see Table 2 for details and accession numbers). One sample was identified only as 'caiman' (YPM 15394) and analyzed as an unknown species. The final source of samples used in this study was wildlife products confiscated by the US Fish and Wildlife Service as illegal imports and donated to this project as research material. Aside from originating in Africa, the specimens (five crocodile skin handbags) contained no additional information on location or species. These unidentified samples were included in the study to test the ability of standard barcode primers to amplify

highly degraded material. Field-collected tissue samples were stored in 10% buffered EDTA-DMSO and kept at room temperature for up to several months before being stored at -20°C ; blood samples were applied directly to Whatman filter paper (Florham Park, NJ), then dried and stored at room temperature for nearly 10 years.

DNA extraction, amplification and sequencing

Samples included in the study had been stored as preserved fresh tissue, dried tissue, blood dried in buffer, blood dried on filter paper, and processed skin products. DNA was extracted from tissue and blood using DNEasy kits (Qia-gen) in a pre-PCR laboratory to prevent contamination. Extractions followed the manufacturer's protocol for buffered animal tissues but were modified slightly for extraction of DNA from dried blood or older tissue to ensure maximum yield from low-quality samples. Modifications included incubating the lysed tissue at 65°C for 15 min after adding AL buffer and incubating again at 4°C for 1 h after adding ethanol. To maximize final yield of genomic DNA, 75 μl of AE buffer, preheated to 70°C , was added and left to incubate for 45 min before centrifuging and collecting flow-through. Because the processed crocodile leather products were the most difficult to extract, we assessed their DNA yield using a NanoDrop ND-1000 spectrophotometer (Thermo Fisher Scientific).

Sequence data were generated using one of three universal *COX1* primer "cocktails" (Table 3; Ivanova et al. 2007). Primer selection for each taxon was made through polymerase chain reaction (PCR) optimization of cocktails 'COI-1,' 'COI-2' and 'COI-3' and one individual primer (VF1d_t1/VR1d_t1) using representative samples from each genus of artiodactyl, primate and crocodilian included in the study, including positive control samples previously sequenced at other gene regions. Table 3 summarizes the taxon-specific primer combinations. PCR was performed in Mastercycler EP gradient S thermocyclers (Eppendorf) in a 25 μl reaction volume containing 1.0 μl genomic DNA (~ 25 ng/ μl), $1\times$ PCR Buffer (Fisher Scientific), 0.24 μM dNTPs, 15 ng BSA, 1 U Taq polymerase (Fisher Scientific), and 0.4 μM of each forward and reverse primer or primer cocktail (Integrated DNA Technologies). Failed amplifications were repeated under the same conditions with 2 μl of genomic DNA. Generally, the optimized PCR thermal cycling profile was a step-up protocol of 94°C for 3 min, 5 cycles of 94°C for 30 s, 51.1°C for 40 s and 72°C for 1 min, followed by 30 cycles of 94°C for 30 s, 56.9°C for 40 s and 72°C for 1 min, with a final extension at 72°C for 10 min. The COI-1 primer cocktail required a different annealing temperature, (54.2°C) for the first 5 cycles. PCR products were visualized by agarose gel electrophoresis, and the cocktail or primer pair yielding the brightest, thinnest band

Table 2 Accession and locality information for museum and US Fish and Wildlife crocodilian specimens sequenced for *COXI* barcoding

Study ID	Institution accession # ^a	Species	Country	Locality	Collection year	Sex
Caiman1	YPM 15394	Caiman	Bolivia	Tarija Dept	1986	
C. c. chiapasius3	YPM 15709	<i>Caiman crocodilus chiapasius</i>	Costa Rica	Puntarenas Province	1993	M
C. c. chiapasius4	YPM 15713	<i>Caiman crocodilus chiapasius</i>	Costa Rica	Puntarenas Province	1993	M
C. c. chiapasius16	YPM 15741	<i>Caiman crocodilus chiapasius</i>	Costa Rica	Alajuela Province	1993	
C. c. chiapasius17	YPM 15743	<i>Caiman crocodilus chiapasius</i>	Costa Rica	Alajuela Province	1993	
C. yacare7	YPM 15402	<i>Caiman crocodilus yacare</i>	Bolivia	SantaCruz Dept	1989	
C. yacare8	YPM 15401	<i>Caiman crocodilus yacare</i> ^b	Bolivia	El Beni Dept	1986	
C. yacare15	YPM 15683	<i>Caiman crocodilus yacare</i>	Brazil	Mato Grosso State	1987	
C. yacare32	YPM 15570	<i>Caiman crocodilus yacare</i>	Brazil	Mato Grosso Estate	1987	F
C. yacare33	YPM 15410	<i>Caiman crocodilus yacare</i>	Bolivia	Pando Dept	1986	
C. yacare34	YPM 15775	<i>Caiman crocodilus yacare</i> ^b	Paraguay	Misiones Dept	1986–1987	
C. yacare35	YPM 15669	<i>Caiman crocodilus yacare</i> ^b	Brazil	Mato Grosso State	1986–1987	
C.c. crocodilus × yacare2	YPM 15547	<i>C.c.crocodilus</i> × <i>yacare</i>	Brazil	Amazonas State	1987	F
C. latirostris5	YPM 15754	<i>Caiman latirostris</i>	Paraguay	Presidente Hayes Dept	1986–1987	
C. latirostris6	YPM 15755	<i>Caiman latirostris</i>	Paraguay	Neembucu Dept	1986–1987	
C. latirostris18	YPM 15392	<i>Caiman latirostris</i>	Bolivia	Tarija Dept	1986	
C. latirostris19	YPM 15554	<i>Caiman latirostris</i> ^b	Brazil	Espirito Santo State	1989	
C. latirostris20	YPM 15551	<i>Caiman latirostris</i> ^b	Brazil	Espirito Santo State	1989	
C. latirostris21	YPM 15548	<i>Caiman latirostris</i>	Brazil	Mato Grosso du Sol	1987	M
C. latirostris22	YPM 15754	<i>Caiman latirostris</i>	Paraguay	Presidente Hayes Dept	1986–1987	
C. latirostris23	YPM 15393	<i>Caiman latirostris</i>	Bolivia	Tarija Dept	1986	
M. niger9	YPM 15833	<i>Melanosuchus niger</i>	Peru	Madre de Dios Dept	1984	
M. niger10	YPM 15834	<i>Melanosuchus niger</i>	Peru	Madre de Dios Dept	1984	
M. niger24	YPM 15695	<i>Melanosuchus niger</i>	Peru	Madre de Dios Dept	1987	F
M. niger25	YPM 15832	<i>Melanosuchus niger</i> ^c	Peru	Madre de Dios Dept	1984	
M. niger26	YPM 15835	<i>Melanosuchus niger</i>	Peru	Madre de Dios Dept	1984	
P. palpebrosus11	YPM 15703	<i>Paleosuchus palpebrosus</i>	Brazil	Mato Grosso State	1987	
P. palpebrosus12	YPM 15702	<i>Paleosuchus palpebrosus</i> ^b	Brazil	Rondonia State	1987	
P. palpebrosus28	YPM 15697	<i>Paleosuchus palpebrosus</i> ^b	Brazil	Rondonia State	1987	
P. palpebrosus29	YPM 15704	<i>Paleosuchus palpebrosus</i>	Brazil	Mato Grosso State	1988	
P. trigonatus13	YPM 15705	<i>Paleosuchus trigonatus</i>	Brazil	Rodinia State	1987	F
P. trigonatus31	YPM 15700	<i>Paleosuchus trigonatus</i>	Brazil	Rodinia State	1987	
P. trigonatus14	YPM 15699	<i>Paleosuchus trigonatus</i>	Brazil	Rodinia State	1987	
C. acutusM7	AMNH R100634	<i>Crocodylus acutus</i>	Mexico	Oaxaca	1967	
O. tetraspisM11	AMNH R75421	<i>Osteolaemus tetraspis</i>	Cameroon	Unknown	Unknown	
1USFWS	USFWS PB800	African crocodile1 ^d	Africa	Unknown	Unknown	
2USFWS	USFWS C5236	African crocodile2 ^b	Africa	Unknown	Unknown	
3USFWS	USFWS C7848	African crocodile3 ^b	Africa	Unknown	Unknown	
4USFWS	USFWS C6688	African crocodile4 ^b	Africa	Unknown	Unknown	
5USFWS	USFWS C3508	African crocodile5 ^b	Africa	Unknown	Unknown	

^a Yale Peabody Museum (YPM); American Museum of Natural History (AMNH); US Fish & Wildlife Service National Wildlife Property Repository (USFWS)

^b Unsuccessfully sequenced

^c Determined to be *C. yacare*

^d Determined to be *C. niloticus*

Table 3 Primers, primer cocktails and universal M13 tails used to sequence all samples included in this study. Table and primer information modified from Ivanova et al. (2007). Position of M13 tail indicated by [M13F] or [M13R]

Name	Ratio	Cocktail name/Primer sequence 5'-3'
COI-1 duikers (<i>Cephalophus</i> spp.), red river hog (<i>Potamochoerus porcus</i>), gray-cheeked mangabey (<i>Lophocebus albigena</i>)		
FF2d	1	TTCTCCACCAACCACAARGAYATYGG
FR1d	1	CACCTCAGGGTGTCCGAARAAYCARAA
COI-2 guenons (<i>Cercopithecus</i> spp.), in bold; bongo (<i>Tragelaphus eurycerus</i>)		
LepF1_t1	1	[M13F]ATTCAACCAATCATAAAGATATTGG
VF1_t1	1	[M13F]TCTCAACCAACCACAAAGACATTGG
VF1d_t1	1	[M13F]TCTCAACCAACCACAARGAYATYGG
VF1i_t1	3	[M13F]TCTCAACCAACCAIAAIGAIATIGG
LepRI_t1	1	[M13R]TAAACTTCTGGATGTCCAAAAAATCA
VR1d_t1	1	[M13R]TAGACTTCTGGGTGGCCRAARAAYCA
VR1_t1	1	[M13R]TAGACTTCTGGGTGGCCAAAGAATCA
VR1i_t1	3	[M13R]TAGACTTCTGGGTGCCIAAIAAICA
COI-3 (crocodilians)		
VF2_t1	1	[M13F]CAACCAACCACAAAGACATTGGCAC
FishF2_t1	1	[M13F]CGACTAATCATAAAGATATCGGCAC
FishR2_t1	1	[M13R]ACTTCAGGGTGACCGAAGAATCAGAA
FR1d_t1	1	[M13R]ACCTCAGGGTGTCCGAARAAYCARAA
M13F (–21)		TGTAAACGACGGCCAGT
M13R (–27)		CAGGAAACAGCTATGAC

was chosen for each taxon. PCR products were purified with Ampure magnetic beads (Agencourt Bioscience) on a Biomek FX robotic platform (Beckman Coulter Inc.). Primer cocktails COI-2 and COI-3 included a modified forward and reverse universal M13 tag (Ivanova et al. 2007). A single primer pair, M13(–21) and M13(–27), was used for bidirectional sequencing reactions of PCR products generated from these two cocktails using BigDye v1.1 chemistry (Applied Biosystems Inc.) on a 3730xl DNA Analyzer (Applied Biosystems, Inc.). Forward and reverse sequences were assembled and edited in Sequencher 4.6 (Gene Codes Corp.) and verified by eye. Contig sequences were aligned using ClustalW (Thompson et al. 1994) as implemented in MEGA 4.0 (Tamura et al. 2007).

Sequence analysis

Unidentified crocodilian and ungulate samples were sequenced but then set apart from the remaining data set. All sequences obtained from species identified through museum collections or by independent morphological evaluation were grouped into their respective taxa and examined for diagnostic molecular characters that could be used as identifiers in future studies. Because species misidentifications are possible in field studies, as well as in the accession of voucher museum specimens, we examined both field and museum samples for obvious errors in species designation. Samples with possible mistaken identities were compared with all other known species to determine if a match could be made based on diagnostic characters.

Because of small sample sizes for several taxa and the associated problems of overestimating diagnostic character sites (Brower 1999; Davis and Nixon 1992), we also confirmed misidentified specimens by means of their placement on phylogenetic trees. We used maximum likelihood (ML) phylogenetic inference as implemented in RAxML 7.0.4 (Stamatakis 2006) using the general time-reversible (GTR) substitution model (Lanave et al. 1984; Rodriguez et al. 1990) with rate heterogeneity parameters modeled by the Gamma (Γ) distribution and four rate categories (Yang 1994). Node support was evaluated with 100 rapid bootstrap replicates (Stamatakis et al. 2008). Outgroups included the American alligator (*Alligator mississippiensis*) for both Crocodylidae and Alligatoridae, the bongo (*Tragelaphus eurycerus eurycerus*) for the ungulate phylogeny, and the rhesus macaque (*Macaca mulatta*) for primates. Resulting trees were inspected for monophyletic groupings and for phylogenetic support values of nodes subtending possibly misidentified samples. Mistakes that could unambiguously be attributed to a species based on matching of diagnostic characters and phylogenetic placement were included in their respective taxonomic group for subsequent analyses.

Sequence variability in *COXI* was evaluated at three hierarchical levels: among conspecifics, among congeneric species and among genera within each of the three orders included in this study. The bongo (*T. eurycerus eurycerus*) and the American crocodile (*Crocodylus acutus*), each represented by only one sample, were excluded from these analyses. Intraspecific comparisons quantified nucleotide diversity for each species based on average nucleotide

substitutions per site (transitions + transversions) using a Tamura-Nei model with pairwise sequence comparisons (Tamura and Nei 1993). To compare congeneric species, we identified fixed nucleotide character differences for every species pair within each represented genus and calculated species divergence using the same Tamura-Nei substitution model. This same approach was used to compare average genus-level nucleotide divergence among orders. Positions that contained a fixed character state among all individuals within a species and an alternate but fixed state at the homologous site in the second species were considered diagnostic at the species level. We used MEGA for genetic diversity and divergence calculations and for visual examination of homologous character state positions among congeneric species. We assessed species monophyly and divergence among sister taxa by observing the resulting ML trees and node support values.

Treating the unknown ungulate, caiman and crocodile samples as a test case for species identification, we compared their sequences against diagnostic characters from our set of known species and included these samples in a reanalysis of ML phylograms. Unidentified samples were assigned to described species based on three criteria. First, an unidentified sample had to be included within a monophyletic group with bootstrap support exceeding 95%. Second, its sequence had to be at least 98% similar to the most common haplotype from a described taxon. Third, its inclusion into a particular monophyletic group had to preserve diagnostic character sites previously identified in that group. We considered these criteria sufficiently conservative to prevent type I errors (incorrectly attributing a sample to the wrong species).

Results

Primer selection and sequencing of degraded tissue and blood

The universal primers of Ivanova et al. (2007) reliably sequenced a 645 bp fragment of *COX1* mtDNA for all mammal and reptile species included in this study. Based on gel visualization, the COI-I cocktail worked optimally for duikers (*Cephalophus* spp.), the red river hog (*Potamochoerus porcus*) and the gray-cheeked mangabey (*Lophocebus albigena*). COI-2 worked best for the bongo, while a single primer pair from this cocktail [V(F,R)1d_t1] was optimal for the guenons (*Cercopithecus* spp.). Cocktail COI-3 worked for all species of crocodilians (Table 3).

Because monitoring of the bushmeat trade using DNA barcoding identification will often require amplification of degraded tissue samples, we evaluated the success of universal *COX1* primers on samples varying widely in age and

curation method. The crocodile skin products proved difficult, with widely varying but generally low average DNA concentrations (13.5 ng/μl, SD = 13.9) and low purity (average 260:280 nm = 1.46, SD = 0.34). We were able to extract template DNA from two of the five leather products but only one of these produced a bidirectional *COX1* sequence (DNA yield = 16.35 ng/μl). We were able to obtain high-quality sequence data from a total of 179 of 204 samples tested (87.7%; Table 1). Of 43 samples that originally failed to sequence, seven were successfully sequenced following a second round of PCR amplification and 11 others produced sequences after a re-extraction of genomic DNA. Of the 25 samples that ultimately failed to yield sequences, nine produced visible PCR bands of the approximate molecular weight of the *COX1* fragment, but failed to sequence even after a second round of extraction and amplification. Final clean trace files of all sequenced samples (with the exception of six previously unidentified specimens, Table 1) were contributed to the BOLD database (Ratnasingham and Hebert 2007) at <http://www.barcodinglife.org> and sequences deposited on NCBI GenBank under accession numbers: GQ144467–GQ144639.

Correction of misidentified specimens

Based on diagnostic sites and branch placement on the ML tree, we discovered two discrepancies in species identification. One caiman sample (*M. niger*25) aligned unambiguously with *Caiman yacare* (Fig. 1a), but was labeled as belonging to *Melanosuchus niger*. It is unclear whether the specimen's label was switched in the lab or the collections, misidentified in the field or incorrectly accessioned; the YPM reptile staff is investigating (G.J. Watkins-Colwell, pers. comm.). One ungulate sample collected in the field and labeled as Peter's duiker (*C. callipygus*YF42) was identical to the most common haplotype of the bay duiker (*Cephalophus dorsalis*; Fig. 1b).

Intraspecific sequence variation

Within-species nucleotide diversity (average rate of nucleotide substitutions per site) in the *COX1* gene ranged from 0.0 to 1.92% (Table 5). The greatest amount of variability was observed in the Nile crocodile (*Crocodylus niloticus*). However, recent studies of this species using several mtDNA and nuDNA markers support a species-level division between northwestern and southeastern Africa (Hekkala 2004; Schmitz et al. 2003), though the geographic extent of each clade is not yet resolved. Eight of our Nile crocodile samples were captured in Gabon and, when we analyzed this geographic group separately, exhibited <0.01% nucleotide diversity (Table 5).

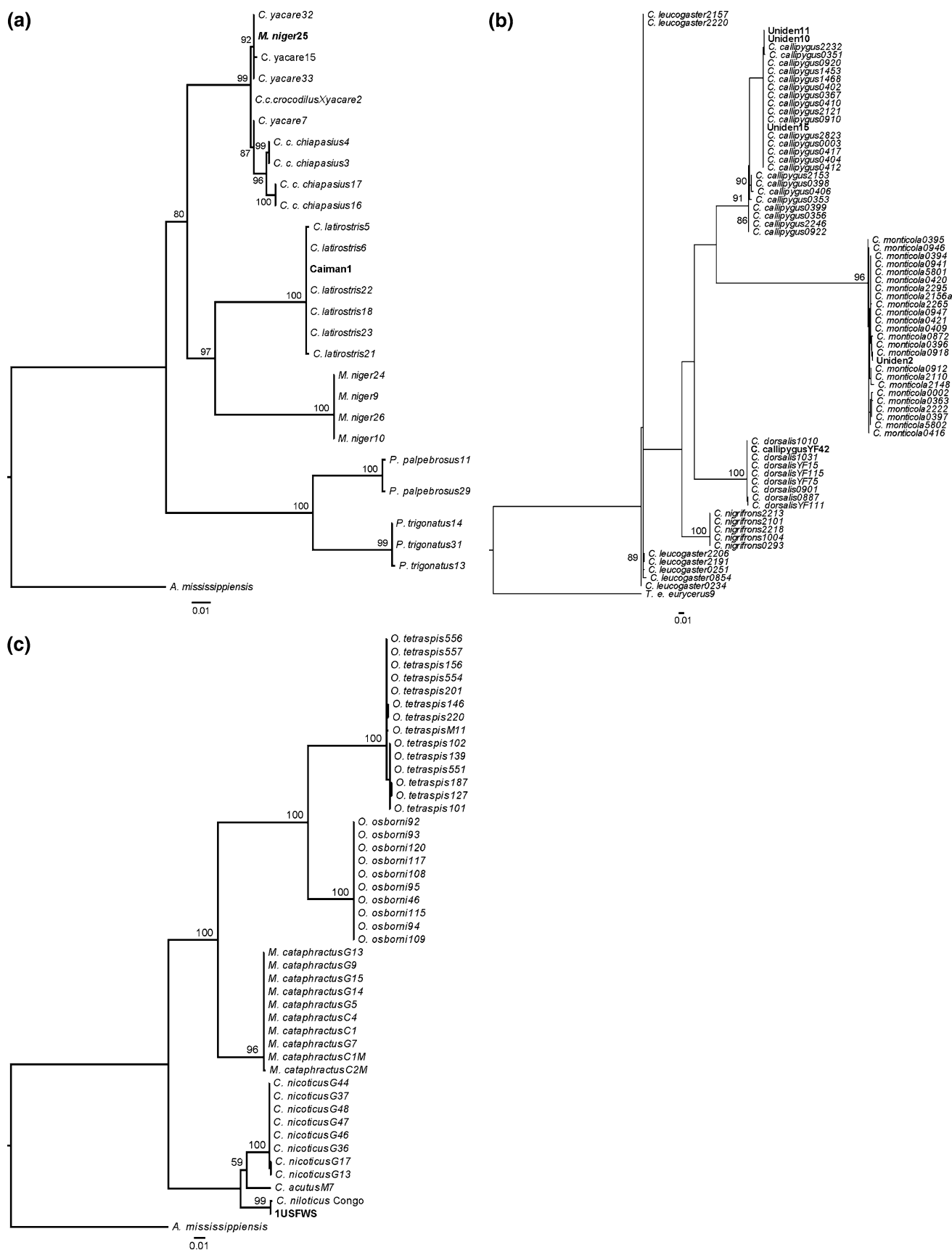


Fig. 1 Maximum likelihood phylograms for **a** South American caimans, **b** African duikers, and **c** African crocodiles. Unknown, blind or misidentified samples are shown in *bold* type. Node support values are based on filtering the best maximum likelihood tree through 100 rapid bootstrap replicate trees. Log-likelihood and alpha shape parameter values, respectively, were **a** −1,949.953015 and 0.199779, **b** −1,888.9429 and 0.02, and **c** −1,965.181261 and 0.248764

We observed the second greatest amount of intraspecific diversity (1.18%) in 26 specimens of Peter's duiker (*Cephalophus callipygus*; Table 5). A subset of eight *C. callipygus* exhibited fixed differences at 13 out of 21 variable sites found within the larger group (Table 5) and formed a paraphyletic assemblage with moderate bootstrap support (<85%, Fig. 1b). When polymorphisms were evaluated separately for the two groups, diversity was 0.41% for the eight individuals and 0.05% in the remaining 18 individuals (Table 5). To reduce the likelihood that we had sequenced a pseudogene in any of the *C. callipygus* samples, we inspected the reading frames and found no premature stop codons. All substitutions were synonymous, thus not resulting in amino acid replacement.

Congeneric species differences

Based on the results of previous phylogenetic (Eaton et al. 2009) and morphological (Brochu 2007) studies, we treated African dwarf crocodiles (genus *Osteolaemus*) from Congo and Gabon as distinct species. We also considered the spectacled caiman (*Caiman crocodilus chiapasius*) and Yacare caiman (*C. yacare*) to be distinct taxa (Busack and Pandya 2001; Vasconcelos et al. 2006) and allowed for the possibility that our Nile crocodile (*Crocodylus niloticus*) samples represent distinct geographic lineages, based on the recent studies noted above.

The number of potentially diagnostic nucleotide characters between congeneric species ranged from 87 sites (13.5% of the sequenced fragment length) between mustached (*Cercopithecus cephus*) and crowned (*C. pogonias*) guenons, to only four characters (0.6% of the *COX1* fragment) between *Caiman yacare* and *Caiman crocodilus chiapasius* (Table 4). Nucleotide divergence values for these species pairs were 0.164 and 0.013, respectively (Table 5). *C. yacare* showed some evidence of paraphyly with *C. c. chiapasius* (bootstrap support = 87%), based on one *C. yacare* sample (*C. yacare*7), while the *C. c. chiapasius* samples themselves grouped more strongly (96%; Fig. 1a).

The average number of diagnostic sites for the 17 congeneric species pairs was 52.3 (8.1% of fragment length; SD = 18.4 sites), corresponding to an average genetic divergence of 0.098 (SD = 0.036). Position and character states of fixed nucleotide sites are presented in Table 4.

Among the three orders compared, average congeneric nucleotide divergence was smallest in the crocodilians (0.065, SD = 0.03; excluding monotypic genera *Melanosuchus* and *Mecistops*) and highest among the three cercopithecoid primate species (0.144, SD = 0.029; excluding monotypic *Lophocebus*). Average pairwise divergence among the five duiker species (0.104, SD = 0.02) was intermediate between primates and crocodilians. Figure 2 demonstrates the range of intraspecific nucleotide diversity found in this study as compared to sequence divergence between congeneric species pairs.

Higher-order comparisons

Comparing sequences between genera revealed average divergence values ranging from 0.084 (*Caiman–Melanosuchus*) to 0.212 (*Cercopithecus–Lophocebus*) within families, to a high of 0.295 between Suidae (*Potamochoerus*) and Bovidae (*Cephalophus*; Table 6). The greatest divergence between Alligatoridae and Crocodylidae was *Paleosuchus* to *Osteolaemus* (0.244; Table 6).

Diagnosis of unknown samples

We determined the species identity of the unknown crocodilian and ungulate samples by matching ostensibly fixed character sites and by placement and likelihood support in the phylogenetic trees. The sequence for the unknown caiman (Caiman1) matched the most common haplotype of *Caiman latirostris* and fell within this monophyletic group with 100% node support (Fig. 1a). The museum sample identified as a hybrid caiman (*C. c. crocodilus* × *yacare*) grouped with *C. c. chiapasius* + *C. yacare*7, but with low bootstrap support (34%; Fig. 1a, node value not shown). This sample, however, matched with *C. yacare* at all diagnostic sites distinguishing the two species (Table 4). The crocodile skin handbag (IUSFWS) matched closely (99.8% identity) with the one Nile crocodile sampled in the Congo. Differentiation of these two samples from Nile crocodiles collected in Gabon received high bootstrap support (99%, Fig. 1c). Comparing these putative groupings revealed 31 potentially fixed character differences and a sequence divergence of 0.053 between Nile crocodile clades (Table 4, 5). One unidentified ungulate sample (Uniden2) matched the most common *Cephalophus monticola* haplotype, while the remaining three (Uniden10, 11, 15) matched the most common *C. callipygus* haplotype. Phylogenetic placement of all four unknown ungulate samples was unambiguous (Fig. 1b). No unknown or misidentified primate samples were detected and therefore the phylogeny is not shown.

for details on taxa and sample numbers and localities). Ambiguity codes denote that this position for a given species is not fixed and that the homologous position for another taxon is diagnostic

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Table 5 Inter- and intraspecific nucleotide differences in the *COXI* gene for three orders of tropical mammals and reptiles. The lower triangular matrix quantifies the number of fixed, diagnostic nucleotide positions between species pairs within each order; values in the upper

matrix represent pairwise nucleotide divergence (Tamura-Nei) across the 645 bp fragment of *COXI*. Diagonal values (in **bold**) are the average number of base substitutions per site ($\times 100$) for intraspecific comparisons

Crocodilia		1	2	3	4	5	6	7	8	9	10	11
1.	<i>M. niger</i> (4)	0.000	0.132	0.129	0.083	0.082	0.086	0.229	0.222	0.212	0.243	0.232
2.	<i>P. palpebrosus</i> (2)	73	0.310	0.066	0.120	0.110	0.118	0.229	0.214	0.213	0.230	0.254
3.	<i>P. trigonatus</i> (3)	74	37	0.100	0.133	0.110	0.119	0.217	0.212	0.208	0.235	0.256
4.	<i>C. latirostris</i> (6)	49	66	75	0.100	0.078	0.082	0.225	0.208	0.211	0.235	0.247
5.	<i>C. yacare</i> (5)	48	62	66	45	0.230	0.013	0.212	0.217	0.208	0.235	0.246
6.	<i>C. c. chiapasius</i> (4)	50	64	67	46	4	0.420	0.225	0.227	0.216	0.244	0.249
7.	<i>M. cataphractus</i> (10)	126	125	121	124	122	126	0.030	0.136	0.136	0.135	0.152
8.	<i>C. niloticus</i> -Gabon (8) ^a	123	118	119	116	123	128	79	0.070	0.053	0.173	0.182
9.	<i>C. niloticus</i> -Congo (2) ^a	114	113	112	114	113	118	76	31	0.160	0.178	0.205
10.	<i>O. osborni</i> (10)	131	125	128	128	129	134	77	96	95	0.000	0.098
11.	<i>O. tetraspis</i> (14)	131	139	111	138	136	139	85	100	106	57	0.280
Artiodactyla		12	13	14	15	16	17	18	19			
12.	<i>C. nigrifrons</i> (5)	0.00	0.103	0.072	0.131	0.082	0.082	0.085	0.327			
13.	<i>C. dorsalis</i> (9)	61	0.06	0.091	0.130	0.097	0.098	0.098	0.289			
14.	<i>C. leucogaster</i> (7)	41	51	0.34	0.124	0.093	0.092	0.099	0.290			
15.	<i>C. monticola</i> (24)	74	69	66	0.53	0.111	0.109	0.119	0.305			
16.	<i>C. callipygus</i> (23)	44	46	46	54	1.18	–	–	0.264			
17.	<i>C. callipygus</i> grp1 (15)	50	56	52	58	–	0.05	0.024	0.318			
18.	<i>C. callipygus</i> grp2 (8)	49	55	54	59	–	13	0.41	0.328			
19.	<i>P. porcus</i> (7)	140	128	125	120	124	132	134	0.13			
Primates		20	21	22	23							
20.	<i>C. cephus</i> (7)	0.150	0.111	0.164	0.240							
21.	<i>C. nictitans</i> (7)	62	0.540	0.157	0.203							
22.	<i>C. pogonias</i> (6)	87	72	0.340	0.194							
23.	<i>L. albigena</i> (8)	118	99	102	0.110							

^a Analyzed as a single species, intraspecific variability of *C. niloticus* was 1.92%

12S rRNA gene for Nile crocodiles from Congo and Gabon (Eaton, unpublished data) to compare with the lineages described by Schmitz et al. (2003) and Hekkala (2004). We found that Gabon crocodiles grouped strongly (ML bootstrap support = 96%) with the assemblage including South Africa, Madagascar, Kenya, Zimbabwe, Sudan and Egypt (GenBank accession numbers: AY195943, AY195945, AY195946, AY195950–AY195955). The Congo specimen and handbag aligned with samples from Senegal, Chad, the Gambia and Mauritania (AY195944, AY195947–AY195949, AY195956, AY195957) with similar bootstrap support (96%). Although it is not possible to compare Nile crocodile *COXI* barcodes directly to clades produced by the 12S rRNA gene, we infer that the differentiation of barcode samples from Congo and Gabon corroborate the earlier studies and represent a geographic division between the former conspecific crocodiles.

Peter's duiker (*Cephalophus callipygus*) was the second species with relatively high levels of intraspecific diversity. A subgroup of eight individuals displayed putative fixed characters at more than half of the variable sites for this species and formed a paraphyletic assemblage (Fig. 1b). Although NuMts (mitochondrial pseudogenes in the nucleus) are not easy to diagnose in cases of noncoding DNA (Kolokotronis et al. 2007), they often contain premature stop codons and frame-shift mutations (e.g. Chung and Steiper 2008; Lemos et al. 1999). Our sequence data contained neither, suggesting NuMts are unlikely to explain the observed sequence variation and phylogenetic pattern. In an extensive DNA barcode survey of birds, Kerr et al. (2007) were able to detect a low prevalence of pseudogenes due to their generally reduced size (100–200 bp) and, presumably, disrupted reading frames. Hybridization with another duiker species is also an unlikely explanation for

Fig. 2 Frequency histogram of intraspecific *COXI* sequence diversity (“species”) compared to nucleotide divergence between pairs of congeneric sister species analyzed in this study. Only sister species *Caiman yacare* and *C. crocodilus chiapasius* had a divergence value overlapping the range of intraspecific nucleotide polymorphisms. The taxonomy of these two species is still being investigated

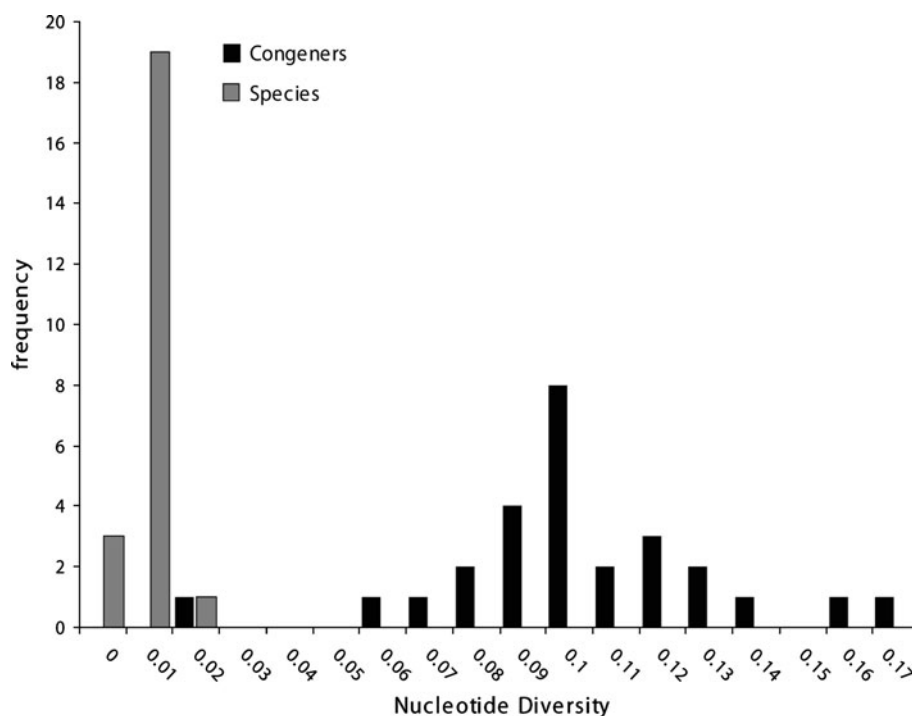


Table 6 Average nucleotide divergence (Tamura-Nei) between genera within orders for a 645 bp fragment of *COXI* mtDNA

Order (Family)	Genera compared	Divergence
Artiodactyla	<i>Cephalophus</i> – <i>Potamochoerus</i>	0.295
Primates	<i>Cercopithecus</i> – <i>Lophocebus</i>	0.212
Crocodilia	<i>Caiman</i> – <i>Crocodylus</i>	0.214
	<i>Caiman</i> – <i>Mecistops</i>	0.221
	<i>Caiman</i> – <i>Osteolaemus</i>	0.243
	<i>Melanosuchus</i> – <i>Crocodylus</i>	0.217
	<i>Melanosuchus</i> – <i>Mecistops</i>	0.229
	<i>Melanosuchus</i> – <i>Osteolaemus</i>	0.238
	<i>Paleosuchus</i> – <i>Crocodylus</i>	0.212
	<i>Paleosuchus</i> – <i>Mecistops</i>	0.223
	<i>Paleosuchus</i> – <i>Osteolaemus</i>	0.244
(Alligatoridae)	<i>Caiman</i> – <i>Melanosuchus</i>	0.084
	<i>Caiman</i> – <i>Paleosuchus</i>	0.118
	<i>Melanosuchus</i> – <i>Paleosuchus</i>	0.131
(Crocodylidae)	<i>Crocodylus</i> – <i>Osteolaemus</i>	0.184
	<i>Crocodylus</i> – <i>Mecistops</i>	0.136
	<i>Osteolaemus</i> – <i>Mecistops</i>	0.143

these findings, as our sample set included all sympatric duikers found in the Republic of Congo with the exception of the yellow-backed duiker (*C. silvicultor*), a much larger species not likely to interbreed with *C. callipygus*. Our observation of cryptic genetic variation in Peter’s duiker is corroborated by at least one other molecular study which found two major clusters of *C. callipygus* in forests of

central Gabon, although the authors offered no discussion of this observation (van Vliet et al. 2008).

Because inference on divergence was based on a single mitochondrial locus within a sympatric population of *C. callipygus*, we cannot reject the possibility that our observations may conflict with the genome tree or true phylogeny for this species due to introgressive hybridization or incomplete lineage sorting (Funk and Omland 2003). Such concerns are reduced in the case of the Nile crocodile, for which analysis of additional genes (including nuclear loci) corroborate those of *COXI* in refuting an imperfect species taxonomy (Hekkala 2004; Schmitz et al. 2003). However, since our results are reported for allopatric populations of Nile crocodiles, introgression may in fact be an important consideration in future analyses if single-gene phylogenies are constructed from specimens sampled in putative hybrid zones.

Interspecific variation

While a character-based approach has been recommended as an improvement over distance-based thresholds for species identification (Rach et al. 2008; Rubinoff et al. 2006), the interspecific differences we present here are based on both distance metrics and diagnostic nucleotide characters. Mean congeneric nucleotide divergence in our study was 9.8% and the average number of fixed characters was 52.3. Variation between congeneric species pairs substantially exceeded levels of intraspecific polymorphism and fell within the range of genetic divergences

reported for a wide range of vertebrate taxa. Hebert et al. (2003b) found that 93.8% of vertebrate congeners had a sequence divergence between 4 and 32% (mean = 9.6%) for the same region of *COXI* while Johns and Avise (1998) reported average genetic distances for congeneric mammal and reptile species generally exceeded ~3% in the mitochondrial cytochrome *b* gene. One exception observed here was between *Caiman yacare* and *C. crocodilus chiapasius*, whose taxonomy is still under debate, but for which a small number of diagnostic characters ($n = 4$) may reliably identify individuals to species (Table 4).

Identification of unknown and misidentified samples

The primary intent of this work was to evaluate whether *COXI* barcoding would serve as a reliable means to identify wildlife species sampled during bushmeat monitoring or while investigating trade in embargoed wildlife products. Matching homologous diagnostic sites and phylogenetic methods unequivocally assigned all unknown samples to the correct species, including caiman, crocodiles and ungulates. Barcode sequences also identified errors in identification or labeling that occurred during field collection or in museum accessioning.

Conclusions

Our results suggest that with minimal effort and simple refinements to DNA extraction and PCR protocols, accurate barcode sequence data can be obtained from most wildlife products encountered in bushmeat monitoring programs and wildlife investigations. Sequencing shorter barcode fragments should increase the success of working with degraded DNA samples and we have begun work to modify universal primers that will contribute to this effort for a wide range of taxa. Strong phylogenetic support and the high frequency of fixed character states between closely related taxa offer convincing evidence that *COXI* barcoding gene will reliably diagnose many common African and neotropical bushmeat species. Bushmeat monitoring and investigations of wildlife commercialization and trade are likely to benefit from this molecular approach, especially when sampling from semi-processed products (e.g. from urban or import markets), when working with products that are difficult to identify (e.g. bird and reptile eggs, fish species, skins), or when focusing in areas containing morphologically cryptic species. In an effort to reduce inevitable errors while conducting field research and in museum curation, mtDNA barcoding offers a simple, low-cost and accurate method for verifying species identities. Although additional independent lines of evidence are needed to substantiate the levels of divergence observed in Peter's

duiker (*C. callipygus*), barcodes may have highlighted a novel evolutionary lineage worthy of further investigation. Finally, generating a database of barcode sequences for tropical wildlife will offer researchers, conservationists and managers an effective tool for more precisely delineating the extent, range and genetic diversity of species of concern.

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